Role of External Pallidal Segment in Primate Parkinsonism: Comparison of the Effects of 1-Methyl-4-Phenyl-1,2,3,6-Tetrahydropyridine-Induced Parkinsonism and Lesions of the External Pallidal Segment


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These experiments re-examined the notion that reduced activity in the external pallidal segment (GPe) results in the abnormalities of neuronal discharge in the subthalamic nucleus (STN) and the internal pallidal segment (GPi) and in the development of parkinsonian motor signs. Extracellular recording in two rhesus monkeys, which had been rendered parkinsonian by treatment with 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP), revealed that the average neuronal discharge rate decreased in GPe but increased in STN and GPi. After MPTP, neurons in all three nuclei tended to discharge in oscillatory bursts. In addition, GABA release in STN (measured with microdialysis) was reduced, indicative of reduced activity along the GPe-STN pathway. Finally, the concentration of glutamic acid dehydrogenase (GAD; measured with autoimmunoradiography) was increased in GPe and GPi, likely reflecting increased striatal input and increased activity of local axon collaterals, respectively. Surprisingly, GAD protein in STN remained unchanged, indicating that the usual assumption that GAD levels are determined primarily by the overall activity of GABAergic elements may be too simplistic. The results from the MPTP-treated animals were compared with results obtained in a second group of three animals with ibotenic acid lesions of GPe. GPe lesions resulted in increased discharge in STN and GPi, comparable with the changes seen after MPTP but did not induce oscillatory bursting and had no behavioral effects. The results indicate that a mere reduction of GPe activity does not produce parkinsonism. Other changes, such as altered discharge patterns in STN and GPi, may play an important role in the generation of parkinsonism.

Key words: subthalamic nucleus; globus pallidus; microdialysis; lesion; Parkinson’s disease; extracellular recording

Introduction

The basal ganglia participate in larger circuits that also include cortex and thalamus. The striatum is the principal input structure of the basal ganglia, and the internal segment of the globus pallidus (GPi) and the substantia nigra pars reticulata (SNr) are the major output structures, projecting toward thalamus and brainstem. According to conventional anatomical models, basal ganglia input and output structures are linked via a monosynaptic “direct” pathway and a polysynaptic “indirect” pathway that involves the external pallidal segment (GPe) and the subthalamic nucleus (STN). Except for the excitatory glutamatergic efferents from STN, intrinsic connections and basal ganglia output projections are GABAergic and inhibitory (Albin et al., 1989; Alexander and Crutcher, 1990). Dopamine released from terminals of the nigrostriatal projection is thought to modulate basal ganglia activity by inhibiting activity along the indirect pathway and enhancing activity along the direct pathway (Alexander et al., 1990). Given the polarity of connections along direct and indirect pathways, striatal release of dopamine may result in an overall reduction of basal ganglia output.

The same model has been applied to explain aspects of the pathophysiology of parkinsonism. Loss of striatal dopamine is believed to result in increased striatal inhibition of GPe, leading to disinhibition of STN neurons and to increased basal ganglia output from GPi and SNr (Wichmann and DeLong, 2003b). Increased and altered basal ganglia output to the thalamus is thought to disturb cortical processing, which is ultimately responsible for the development of many of the parkinsonian motor signs.

Although generally supported by previous recording and lesioning studies, some of the more recent anatomical and biochemical studies have provided findings that are difficult to reconcile with this model. In particular, these reports have cast doubt on the notion that discharge abnormalities in STN and GPi

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and parkinsonian motor abnormalities are attributable to reduced GPe activity. Thus, comparisons between 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)-intoxicated monkeys or parkinsonian patients and their respective controls failed to reveal a significant parkinsonism-related decrease in the number of GPe output neurons expressing the mRNA coding for the GABA-synthesizing enzyme glutamic acid decarboxylase (GAD) (Herrero et al., 1993, 1996a,c), a marker presumed to reflect the electrical activity of GABAergic neurons. Furthermore, the expression of mRNA coding for cytochrome oxidase subunit I was significantly increased in parkinsonian monkeys in neurons of STN, GPi, and SNr, confirming that their activity increased after nigrostriatal denervation, but there was no statistically significant change in GPe (Vila et al., 1997).

The aim of this study was to re-examine whether the GPe-STN projection is underactive in parkinsonism, and whether reduction of neuronal activity in GPe leads to parkinsonian motor signs and electrophysiological abnormalities in STN and GPi similar to those found in parkinsonism (for related experiments in rodents, see Hassani et al., 1996). We measured MPTP-induced changes in neuronal activity and GAD expression in GPe, STN, and GPi, as well as GABA release in the STN. These data were compared with data obtained in monkeys with GPe lesions.

Materials and Methods

General methods

Five animals were used for these studies. All animals underwent initial electrophysiological recording of the activity of STN, GPi, and GPe. In this phase of the experiments, the animals' behavior was also quantified using a battery of behavioral assessment methods.

Two of the animals (monkeys H and I) received intracarotid injections of MPTP to induce hemiparkinsonism. Before and after MPTP treatment, GABA levels in the STN were measured with microdialysis methods as an indication of GABA release from terminals of the GPe-STN projection. In addition, post-MPTP recording studies were performed.

The remaining three animals (monkeys F, K, and X) received lesions of GPe followed by additional recording and behavioral observations. At the termination of the experiments, histological studies were performed in all five animals to verify the targets of electrode penetrations and to delineate the extent of dopaminergic and pallidal lesions.

Animals

The five rhesus monkeys (Macaca mulatta; 4–5 kg) used for these studies were housed under conditions of protected contact housing with ad libitum access to standard primate chow and water. Before any of the other procedures, the animals were trained to sit in a primate chair, to adapt to the laboratory, and to permit handling by the experimenter. During injection, recording, and microdialysis experiments, the animals were awake. All experimental protocols were performed in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and the United States Public Health Service Policy on Humane Care and Use of Laboratory Animals (amended 2002). All experiments were approved by the Institutional Animal Care and Use Committee of the Emory University Institutional Biosafety Committee.

Surgical procedure

After completion of behavioral training, metal chambers for chronic recording (inner diameter, 16 mm) were stereotactically positioned over a trephine hole under aseptic conditions and isoflurane anesthesia (1–3%). A chamber directed at the pallidum was placed at an angle of 50° from the vertical in the coronal plane [anterior (A), 12; lateral (L), 10; dorsal (D), 5 Horsley–Clark plane (HcO)], and a chamber aimed at the STN was placed at an angle of 36° from the vertical in the sagittal plane (A, 10; L, 6; D, 2 HcO). The chambers were affixed to the skull with dental acrylic. Metal head holders were also embedded into the acrylic cap to permit head stabilization during the recording and microdialysis procedures.

Electrophysiology experiments

The neuronal activity in GPe, GPi, and STN was recorded extracellularly with tungsten microelectrodes (impedance, 0.5–1.0 MΩ at 1 kHz; Fredrick Haer, Bowdoinham, ME). A microdrive (MO-95B; Narishige, Tokyo, Japan) was used to lower the microelectrodes through the dura into the brain, using a 20 gauge guide tube. The electrical signal was amplified (DAM-80 amplifier; World Precision Instruments, Sarasota, FL), filtered (400–10,000 Hz; Krohn-Hite, Brockton, MA), displayed on a digital oscilloscope (DL1540; Yokogawa, Tokyo, Japan), and made audible via an audio amplifier. A tape record of the signal was kept using a video recording adapter (model 3000A; Vetter, Rebersburg, PA). Neurons in the basal ganglia were easily identified by such characteristics as a high-frequency discharge with pauses in GPe, tonic high-frequency discharge in GPi, and a tonic and regular activity in an area of high background activity in the STN.

Because none of the GPe lesions was complete, the possibility exists that the lesion effects may have been specific for a particular topographic region within the GPe (for instance, the motor or associative territories). To detect the effects of GPe inactivation, it would thus be desirable to restrict recording to functionally related areas of GPe and STN. With the lateral coronal approach to the globus pallidus chosen for this study, functionally related regions of GPe and GPi are (approximately) aligned along a given electrode track. Thus, by restricting GPe recordings to those cells that followed an electrophysiologically silent (lesioned) portion of GPe, we restricted recording in GPi to cells that are (presumably) functionally related to the lesioned portion of GPe. It was not possible to restrict the analysis in a similar manner in the STN.

Microdialysis

 Probe assembly and calibration. Dialysis probes were made with a dual silica tube, parallel flow design (Olson and Justice, 1993). The shaft of the probe was 15 cm long and was made of 27 gauge stainless-steel tubing (Small Parts, Miami, FL). The inflow and outflow tubes were inserted into the 27 gauge stainless-steel tubing and consisted of two pieces of fused silica [inner diameter (i.d.), 0.04 mm; outer diameter (o.d.), 0.1 mm; Polymicro Technologies, Phoenix, AZ]. The inflow tube extended beyond the distal end of the outflow tube by the length of the active dialysis area of the membrane. A short piece of hollow dialysis membrane tubing (i.d., 0.2 mm; cutoff molecular weight, 13,000; Polymicro Technologies) was placed over the ends of the inflow and outflow silica tubes. Epoxy glue (two-ton clear epoxy; Devcon, Chicago, IL) was used to seal the open end of the microdialysis membrane and the junctions between the microdialysis membrane and the inflow and outflow silica tubes. The assembled dialysis membrane and the inflow and outflow silica tubes were glued to the 27 gauge stainless-steel tubing with white epoxy (Devcon). The exposed surface of the hollow dialysis membrane tubing of the probe was 2 mm. The inflow line of the microdialysis system was connected to Teflon tubing (o.d., 0.65 mm; i.d., 0.12 mm; Bioanalytical Systems, West Lafayette, IN) via a liquid switch syringe selector (CMA Microdialysis, Solna, Sweden) to a 1ml syringe (Bioanalytical Systems) and perfused with artificial CSF (aCSF; in mm: 145 NaCl, 2.8 KCl, 1.2 MgCl₂, 1.2 CaCl₂, pH 7.2–7.4) at a flow rate of 2 µl/min with a microdialysis syringe pump (Bioanalytical Systems). For a 20 min period in each experiment (see below), the system was perfused with aCSF containing 80 mM K⁺. In these experiments, the concentration of NaCl was lowered to maintain osmolality (in mm: 67.8 NaCl, 80 KCl, 1.2 MgCl₂, and 1.2 CaCl₂).

The dead volume of connecting tubes and the probes themselves were calculated to synchronize microdialysate collection with the timing of switches to aCSF exposure. The in vitro relative recovery was calculated by determining the ratio of GABA peak height measured in an equivalent volume of aCSF containing a known amount of GABA. The recovery of the probes ranged from 15 to 20% at flow rate of 2 µl/min by using the aCSF solution.

Microdialysis protocol. Before use, the probes were placed into aCSF and were thoroughly flushed with aCSF at a rate of 2 µl/min for 20 min.
The probes were then slowly inserted into the STN using the microdrive and guide cannula described above. The exact target coordinates were determined by the results of the previous electrophysiological recording experiments. Before sample collections, the probes were perfused with normal aCSF for 2 hr to allow stabilization of the basal GABA concentration before the start of sample collection. Pilot experiments with the techniques used here have shown that the measured GABA levels stabilized 90–120 min after insertion of the microdialysis probes into the brain. In each experiment, eight dialysis samples were collected (20 min; 40 μL/sample) in plastic microcentrifuge tubes. The first five samples were collected without additional intervention, but the perfusion media was changed for the duration of sample 6 to one containing 80 mM K+ (see above) using a liquid switch syringe selector (CMA Microdialysis). The potassium stimulation of the tissue was used primarily to assess the neuronal origin of GABA.

For the last two samples of the protocol, the probe was again perfused with standard aCSF solution. All samples were immediately placed on dry ice and later stored at −80°C until the time of analysis. At the termination of each microdialysis session, the probe was removed, the chamber was cleaned, and the animal was put back in its home cage. The same animal underwent multiple microdialysis procedures, separated by at least 24 hr. For each microdialysis session, a new track was used to sample undisturbed tissue.

**Administration of MPTP**

After completion of recording and microdialysis procedures in the normal state, monkeys H and I received MPTP by injection into the right carotid artery (0.5 mg/kg per injection; monkey H received two injections 36 d apart; monkey I received one injection), following reported standard protocols (Bankiewicz et al., 1986). Both animals showed clear parkinsonian signs on the side contralateral to the injections (i.e., on the left side of their body) after these injections, including bradykinesia, rigidity, and flexed limb posture. The post-MPTP experiments started 2 months after the last MPTP injection.

**GPe lesioning procedure**

These studies were performed in monkeys F, K, and X. The animals received multiple small (0.2–0.5 μL, 0.2 μL/min) injections of the flurbiprofen excitoxin ibotenic acid (10 mg/ml, dissolved in bidistilled water on ice, final pH 7.4; Sigma, St. Louis, MO) into GPe. The injections were performed with a combined injection-recording device, which permits electrophysiological recording before and during injections to verify injection location. The device consists of a 30 gauge injection cannula through which a Teflon-coated tungsten recording wire (0.05 mm bare diameter, 0.076 mm coated diameter; AM Systems, Carlsborg, WV) was threaded, the tip of which protrudes 0.2–0.7 mm from the cannula. The device was passed through the dura through a guide tube. Gray and white matter areas of the brain were easily identified with this system, facilitating the identification of injection targets. The injections were delivered over several days. The volume of the individual injections was kept to a minimum to limit the spread of the toxin. Between drug injections, the amount of damage produced by the toxin was assessed by additional electrophysiological mapping. The series of injections was terminated once the accessible GPe was electrophysiologically silent. Monkey F received a total of 115 μg of ibotenic acid, whereas monkey K received 33 μg and monkey X received 29 μg.

**Behavioral assessment method**

Behavioral changes associated with GPe inactivation, as well as parkinsonian signs associated with MPTP treatment, were documented through observations of spontaneous cage behavior. For quantification, one of the investigators (J.S. for the MPTP-treated animals; M.K. for the GPe lesioned animals) used a computer-assisted behavioral scoring system to score the occurrence of spontaneous limb movements in an observation cage over a 20 min period. The system has been used previously and has been validated in the context of other studies (Bergman et al., 1990; Wichmann et al., 2001a). Briefly, each time the animal moved one of its extremities, a computer keyboard key assigned to this extremity was pressed and held down for the duration of the movement. The ratio of arm movements registered on the left side and those registered on the right side was used as an index of arm movements.

We also used an automated activity monitoring system. The observation cage was equipped with eight infrared beams. Four of these beams were arranged in a square design between the back and the front of the cage, and four additional beams, arranged in a similar manner, between the two sides of the cage. An attached computer kept a log of the timing of infrared beam crossings. Off-line, the system then calculated total activity counts. The 20 min activity monitoring periods were also videotaped.

All observations were performed at the same time of day. The behavioral assessment method was used before and twice weekly after completion of the MPTP treatment and of GPe lesions.

**Histology**

At the conclusion of the experiments, the monkeys were killed by induction of deep anesthesia with an overdose of sodium pentobarbital, followed by transcardial perfusion with saline and 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.2. The brains were removed and cryoprotected in a 30% sucrose solution in 0.1 M PB. The fixed brain was sectioned in coronal planes (50 μm). One of every four sections was stained with cresyl violet. In the case of the MPTP-treated animals, the remaining sections were used for double-labeling studies to visualize GAD and tyrosine hydroxylase (TH). In the GPe-lesioned animals, sections were used to label the neuronal marker microtubule-associated protein 2 (MAP2) using standard immunohistochemistry protocols.

For the double-labeling studies, sections were thoroughly washed in 0.1 M Tris-buffered saline (TBS), to remove the cryoprotectant and incubated in 10% normal goat serum with 0.04% Triton X-100 in TBS for 30 min. Thereafter, they were incubated for 72 hr in the same solution containing either a primary antisera or a mixture of primary antibodies. The antibodies used for this study were against TH (1:1000, mouse monoclonal antibody; Chemicon, Temecula, CA) and GAD (1:2000, rabbit polyclonal antibody; Chemicon). Next, the sections were rinsed in 0.1 M TBS and incubated for 2 hr with goat anti-rabbit antibody conjugated to 125I (for GAD) and biotinylated goat anti-mouse antibody (for TH). For enzyme-linked staining of TH-positive cells or GAD, the biotin–avidin complex method was used, whereas 3,3′-diaminobenzidine tetrachloride was used to visualize the final product. For control sections, either one or both of the primary antibodies were omitted. Finally, the sections were rinsed in 0.1 M TBS, mounted on gelatin-coated slides, and air-dried. The slides were placed in x-ray cassettes with calibrated 125I standards (Amersham Biosciences, Piscataway, NJ) and exposed to Hyperfilm3H (Amersham Biosciences) for 2–3 weeks. The films were developed in D-19, fixed in Kodak (Rochester, NY) fixer, and dried.

**Data analysis**

**Analysis of behavioral data.** To analyze the detailed cage observation records, a ratio of the durations was computed (within the 20 min observation period), during which the left and the right arm, or the left and right leg, were used. These left/right (L/R) ratios were used as indicators of parkinsonian hypokinesia (Bergman et al., 1990). The L/R ratios and activity monitoring scores were then plotted over time, comparing the post-MPTP or postlesion data with data obtained in the normal state. Because there were no obvious changes over time in the L/R ratios or activity counts after the MPTP or GPe lesions, the postintervention data were pooled for each monkey and compared with the preintervention data for the respective animal.

**Analysis of electrophysiological data.** Only those cells in which reconstruction of their location, on the basis of stereotactic information, location, and depth data gathered during the actual recordings as well as histological analysis of the cresyl violet stained sections, confirmed that they were within the target structures (GPe, GPi, or STN) were included in the analysis. To generate interspike interval (ISI) information, the tape-recorded activity was passed through a template-matching spike sorter (Alpha-Omega, Nazareth, Israel). The subsequent analysis used the Matlab software environment (MathWorks, Natick, MA). Neuronal data were included in this study only if the isolation of the respective cell could be maintained throughout the record, if at least 1000 action potentials were observed and held down for the duration of the movement. The ratio of arm movements registered on the left side and those registered on the right side was used as an index of arm movements.

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tonics had been recorded, and if the spontaneous activity of the neuron was recorded for at least 45 sec. Most neurons were, in fact, recorded for longer (mean, 339.5 sec). The ISI data were converted into frequencies, and a power spectral analysis of the frequency representation (sampled at 100 Hz) was performed, using Hanning windows and mean detrending of data segments. For statistical analysis, the elements of the ISI data stream were randomly shuffled, and the power spectrum of the resulting (random) data stream was calculated. Both power spectra were smoothed with a 10 point moving average. The mean and SD of the resulting power spectrum of the shuffled data were computed. Peaks in the power spectra of the original data stream were considered significant only if 10 consecutive points of the spectrum were at least 5 SD different from the mean of the shuffled data.

Bursts were detected using an algorithm developed by Legendy and Salzman (1985). The algorithm is based on the Poisson “surprise” method. For this analysis, a surprise value of 10 was used. For each cell, we calculated a burst index (i.e., the ratio between the number of spikes found in bursts and the total number of spikes).

**Chemical analysis.** The concentration of GABA in the dialysate samples was determined by HPLC with electrochemical detection as described previously by Donzanti and Yamamoto (1988). Briefly, GABA was assayed by precolumn derivatization with o-phthalaldehyde. The derivatization reagent was prepared by dissolving 27 mg of o-phthalaldehyde in 1 ml of 100% methanol, 9 ml of 0.1 M sodium tetraborate, pH 9.4, and 10 ml of β-mercaptoethanol. This stock solution was then diluted 1:3 with 0.1 M sodium tetraborate. A 10 ml aliquot of the reagent solution was then added to 20 ml of the dialysate sample. Derivatization was allowed to proceed for 30 sec before injection onto a 3 mm ϕ18 reverse-phase column (100 × 2.0 mm; Phenomenex, Torrance, CA) maintained at 32°C. GABA was eluted with a 0.1 M sodium phosphate mobile phase, pH 6.4, containing 23% methanol and 50 mg/l Na2EDTA. Mobile phase was pumped through the system at a flow rate of 0.3 ml/min. Detection of GABA was at a glassy carbon electrode (6 mm diameter) maintained at −0.7 V by an LC4B amperometric detector (Bioanalytical Systems). The limit of detection for GABA was 8.5 pg/20 ml (3:1 signal-to-noise ratio).

**Analysis of microdialysis data.** The effect of potassium stimulation was clearly visible in sample 6 (the time of stimulation) and sample 7 (wash-out period). In each experiment, the basal level of GABA was calculated as the average of samples 1–5 and 8. The stimulation-evoked GABA overflow was calculated by subtracting the basal level (as defined above) from samples 7 and 8 and adding the resultant measurements. For graphical representation (see Fig. 3 A, B), all values were normalized to the average basal level in the normal state.

**Analysis of double-labeling studies.** Images were collected on a Leitz (Wetzlar, Germany) microscope linked to a microcomputer imaging device (MCID) image analysis system (Imaging Research, St. Catharines, Ontario, Canada). For final output, images were composed using Adobe Photoshop (Adobe Systems, San Jose, CA).

For densitometric measurements of GAD immunoreactivity (GAD-IR), the autoradiograms were analyzed with a video-based MCID image analysis system by relating optical density to calibrated 125I standards. Eight sections from the globus pallidus and six sections from the subthalamic nucleus from each of the monkeys used for the study were analyzed. Data from the untreated side were compared with the data from the lesioned side in unilaterally treated monkeys. GAD-IR was expressed as percentage control.

**Analysis of lesion location in the GPe-lesioned animals.** Microscopic images of cresyl violet and MAP2-stained sections were collected using a Leitz microscope. The lesion extent, as judged by the absence of MAP2 staining, was first drawn on serial images of the cresyl violet-stained sections. The outlines of the basal ganglia nuclei as well as the lesion locations were then transferred into line drawings (see Fig. 5). The approximate anterior coronal coordinate of the planes shown in Figure 5 was estimated on the basis of atlas information (Winters et al., 1969) and the number of the histological slices within the series of sections.

**Statistical analysis.** Statistical comparisons were performed in the Statistical Program for the Social Sciences environment. The neuronal data from the different monkeys within the same experimental group or period (normal, MPTP-lesioned, GPe-lesioned) were pooled after exclusion of intersubject differences with ANOVA. Statistical tests for significance were performed with t tests (with Bonferroni correction for multiple comparisons). A p level < 0.05 was accepted as indicating a significant difference.

**Results**

**Behavioral observation.** Both of the MPTP-treated animals developed marked hemiparkinsonism on the side contralateral to the injections. On the affected (left) side, manipulations of objects was greatly impaired, the animals stopped grooming or feeding themselves with the affected arm, and they held their arm and leg in a flexed position. The whole body posture was also markedly stooped. However, by using the right arm, the animals were able to maintain themselves and, in fact, gained weight in the post-MPTP phase. The global activity of the animals was not markedly impaired, because they developed a prominent tendency to circle toward the hemiparkinsonian side in their cages.

Behavioral assessment with the computer-assisted observation system (Bergman et al., 1980) revealed that spontaneous arm and leg movements became markedly asymmetric because of reduced use of the parkinsonian arm. In monkey 1, the L/R ratio for arm movements decreased from a pre-MPTP value of 0.83 ± 0.15 (n = 4) to a post-MPTP value of 0.08 ± 0.05 (n = 16; p < 0.01). In monkey H, only one pre-MPTP assessment was available, showing an L/R ratio of 0.75. After MPTP, the L/R ratios decreased to 0.32 ± 0.06 (n = 8). Similar results were obtained for leg movements. Throughout the recording and microdialysis sessions, the behavioral state remained stable in both animals. All of these findings are typical responses to intracarotid MPTP treatments.

In the GPe-lesioned animals, parkinsonism or other motor abnormalities were not detected by direct cage observations, activity monitoring, or the computer-assisted behavioral observation system.

**Electrophysiological recordings.** Because most of the observed effects were consistent between animals, we will describe these changes here only as averages across monkeys (Figs. 1, 2). The electrophysiological data for individual animals are listed in Table 1.

**Discharge rates.** MPTP treatment resulted in obvious changes in discharge rates and discharge patterns in GPe, STN, and GPi (Fig. 1, Table 1). In the MPTP-treated state, the average neuronal discharge rate (Fig. 1A) in GPe decreased from a control value of 65.1 ± 2.9/sec (n = 66 neurons; mean ± SEM) to 45.9 ± 2.6/sec (n = 53 neurons; p < 0.001; t test). In contrast, MPTP treatment induced an increase in the average discharge rate of STN neurons from a control value of 25.7 ± 1.2/sec (n = 85) to 36.1 ± 1.6/sec (n = 45; p < 0.005), and in GPi from 65.1 ± 1.8/sec (n = 134) to 80.6 ± 3.1/sec (n = 40; p < 0.001). There were no discernible regional differences in the distribution of GPe, STN, or GPi cells with high or low discharge rates.

Prominent discharge rate abnormalities were also evident in the GPe-lesioned animals. After the lesion, the average rate of discharge in the STN rose from the above-mentioned control of 25.7 ± 1.2 to 36.0 ± 2.2/sec (n = 74; p < 0.01). This was significant in both animals for which STN data were available (Table 1). In GPi, the average rate rose from 65.1 ± 1.8 to 69.3 ± 2.0/sec (n = 137). The mean rate increased in all three animals, but this reached significance only in one (Table 1) and was not significant in the aggregate. There were no discernible regional differences in the distribution in STN or GPi of cells with high or low discharge.
rates. However, as mentioned in Materials and Methods, the recording in GPi was restricted to an area functionally related to the lesioned portion of GPe, whereas no effort was made to restrict the recording to any particular territory within STN. GPe lesions and MPTP-induced depletion of striatal dopamine resulted in similar changes in the overall discharge rates in STN and GPi.

Discharge patterns
In addition to the abnormalities in neuronal discharge rate, there were also obvious changes in discharge patterns in the MPTP-treated animals (Figs. 1, 2). The incidence of burst discharge (Fig. 1B), subjectively noted during the recording sessions and objectively assessed with the burst-detection algorithm, increased in GPe from 0.47 ± 0.06% in the normal state to 1.1 ± 0.2% in the MPTP-treated state (p < 0.05), in STN from 1.28 ± 0.15 to 2.28 ± 0.22% (p < 0.001), and in GPi from 0.32 ± 0.03 to 0.98 ± 0.14% (p < 0.001).

In contrast, the incidence of burst discharges did not change in the GPe-lesioned animals in STN or GPi, either subjectively or objectively. In STN, the postlesion incidence of bursts was 1.08 ± 0.18% (see above for control values), and in GPi, the postlesion incidence of bursts was 0.44 ± 0.02% (Fig. 1B).

Changes in oscillatory discharge were assessed using autocorrelograms and power spectra (as detailed in Materials and Methods). As noted previously, recordings in the MPTP-treated animals revealed that neurons in GPe, STN, and GPi developed a strong tendency to burst in an oscillatory manner in the parkinsonian state. Integration of power spectra in 1–3, 3–8, 8–15, and 15–50 Hz spectral frequency bands revealed that MPTP treatment induced substantial changes in the power spectral content of the neuronal discharge. In all three structures, the 3–8 and 8–15 Hz bands became more prominent (Fig. 2A, Table 1). In STN and GPi, the spectral content in the 1–3 Hz range was also enhanced (data not shown). In GPe, the 1–3 Hz frequency range was less strongly represented. In all structures, the spectral content in the 15–50 Hz range was reduced after MPTP treatment (data not shown). Autocorrelation analyses showed substantial increases in the number of cells with peaks in their autocorrelograms indicative of oscillations in the 3–8 and 8–15 Hz ranges (Fig. 2B). In contrast, neither the power spectral analysis nor the analysis of autocorrelograms revealed any changes in the incidence of oscillatory discharge in STN or GPi in the GPe-lesioned animals (Fig. 2A,B). In fact, the postlesion recordings were virtually identical in this regard compared with those obtained before lesioning. These results indicate that GPe-lesioned and MPTP-treated animals differed substantially with regard to the induction of phasic discharge abnormalities.

Microdialysis experiments
These experiments were aimed at elucidating the effects of dopamine depletion and were, therefore, only performed in monkeys H and I before and after treatment with MPTP (Fig. 3). The basal concentration of GABA in microdialysate samples from the STN in the normal state amounted to 93.8 ± 6.9 pg/20 μl in monkey H and 65.5 ± 7.1 pg/20 μl in monkey I. As shown in Figure 3, A and B, baseline GABA levels were stable throughout the sampling period. Exposure of the tissue to 80 mM K+ resulted in a substantial increase in GABA eflux. The evoked overflow amounted to 3.4 times the basal level of GABA in monkey H and 3.3 times the basal level of GABA in monkey I. Compared with the normal state, the basal level of GABA was reduced by an average of 45.3% in monkey H and 35.2% in monkey I (Fig. 3C). The stimulated overflow was reduced by 52% in monkey H and 47% in monkey I.

Histology
Cresyl violet staining of coronal sections showed gliosis along microelectrode tracts in all of the animals. Although reconstruction of microelectrode tracts aimed at GPe or GPi was straightforward, tracts aimed at the STN were difficult to reconstruct because these tracts were not in the plane of histological sections. All data from penetrations in which placement of the probes were in doubt were excluded from the analysis. In the MPTP-treated animals, immunohistochemistry for TH (Fig. 4A, top) showed that there was a massive loss of dopaminergic terminals in the striatum on the side of the MPTP treatment (right). The loss was most pronounced in the dorsal striatum, whereas the dopaminergic supply to the ventral striatum was relatively preserved.

GAD staining in the MPTP-treated animals revealed that the distribution of GAD-IR was uniform throughout the neuropil of both GPe and GPi in control animals. GAD-IR was also observed in the STN. GAD-IR was increased in the GPe and GPi on the lesioned side in the unilaterally MPTP-lesioned monkeys (Fig. 4B). The densitometric analysis confirmed an increase in GAD-IR in GPe and GPi (Fig. 4C) (p < 0.05). In contrast, GAD-IR was not significantly changed in the STN (Fig. 4C) on the parkinsonian side.

Cresyl violet and MAP2 staining of coronal sections throughout the basal ganglia revealed that all three of the GPe-lesioned animals had substantial lesions covering portions of GPe (Fig. 5). The recording chamber placement in monkey F precluded lesioning of the entire GPe. In this animal, the lesion covered the dorsal (predominantly nonmotor) territory of GPe. In the other two animals, the lesions were more extensive and included nonmotor and motor territories of the nucleus. In each case, the core
of the lesion (as judged by MAP2 staining) was surrounded by a halo of gliosis, which was readily visible on cresyl violet staining. Despite our attempts to limit the lesion to GPe, some of the neighboring portions of striatum and GPi were involved in these (fiber-sparing) lesions.

**Discussion**

Our results corroborate the view that GPe activity is reduced in parkinsonism, and that this reduction may account for the changes in discharge rates in STN and GPi in this disorder. However, a mere reduction in GPe activity does not appear to be sufficient for the development of changes in basal ganglia discharge patterns or motor dysfunction in parkinsonism.

**Global neuronal activity changes in parkinsonism**

The findings of reciprocal discharge rate changes in GPe and GPi, and of increased rates in STN (Miller and DeLong, 1987; Albin et al., 1989; DeLong, 1990; Filion and Tremblay, 1991; Bergman et al., 1994), have given rise to the notion that most parkinsonian abnormalities result from changes in the indirect pathway of the basal ganglia (Albin et al., 1989; DeLong, 1990). Striatal dopamine loss is postulated to result in increased striatal inhibition of GPe, which in turn results in disinhibition of STN and increased basal ganglia output from GPi. This concept is supported by the fact that STN inactivation results in a substantial reduction of the severity of parkinsonian motor signs in monkeys (Bergman et al., 1990; Guridi et al., 1993; Baron et al., 2002) and in human patients (Sellal et al., 1992; Gill and Heywood, 1997; Alvarez et al., 2001).

Our electrophysiological and microdialysis findings confirm the proposed global activity changes in GPe, STN, and GPi in parkinsonism. We found that the average discharge rate of GPe neurons was significantly decreased. As a further indication of reduced activity of GABAergic GPe cells, the release of GABA in the STN, which is derived in large part from GPe inputs, was reduced by ~50%. Measurements of discharge rates in subsequent stages of the indirect pathway (STN and GPi) confirmed that average discharge in these areas is substantially increased in parkinsonism, which is likely the result of reduced inhibition via GPe efferents (see also below).

The interpretation of the GAD protein measurements is less straightforward. The levels of GAD protein or of GAD mRNA are often used as measures of the activity of GABAergic cells or terminals. This explanation can serve to interpret the finding of increased GAD protein levels in GPe, which may result from increased GABAergic inputs from striatum (Robertson et al., 1991).

GAD protein levels in GPi were also increased in the MPTP-treated animals. In general, GAD levels in GPi may reflect the activity of GABAergic afferents from striatum or GPe or of local axon collaterals. However, it is unlikely that activity changes in the two major GABAergic afferent pathways account for the observed change in GAD activity. As mentioned above, based on electrophysiological and microdialysis findings, the activity of GPe neurons and, thus, of the GPe-GPi projection is reduced in parkinsonism. Based on measurements of striatal GAD mRNA levels, it is also known that striatal neurons projecting to GPi are not overactive in MPTP-treated monkeys (Soghomonian and La- prade, 1997). It is therefore most likely that the increased level of GAD protein in GPi represents increased activity of local axon collaterals of GPi neurons and is therefore another indicator that GPi activity is enhanced in MPTP-treated animals. This is also suggested by studies that have demonstrated that GAD mRNA levels in GPi are increased in parkinsonism (Pedneault and Soghomonian, 1994; Soghomonian et al., 1994; Herrero et al., 1996a; Salin et al., 2002; Schneider and Wade, 2003).

GAD protein levels in the STN would be expected to reflect changes in the activity along the GPe-STN pathway. The finding of unchanged GAD levels in the STN in our study mirrors the results from previous studies, which have shown that GAD65 mRNA remains the same after induction of parkinsonism (Soghomonian and Chesselet, 1992; Pedneault and Soghomonian, 1994; Schneider and Wade, 2003), whereas the level of GAD-67 mRNA in GPe was increased (Soghomonian and Chesselet, 1992; Delfs et al., 1995; Herrero et al., 1996a; Schroeder and Schneider, 2001; Schneider and Wade, 2003). It is obvious that these findings do not agree with the electrophysiological recording and microdialysis data that indicate that GPe activity is substantially decreased. Thus, the assumption that measurements of GAD mRNA or GAD protein levels simply reflect the overall activity of GABAergic neurons may not apply in this case. The regulation of GAD mRNA or protein may be influenced by factors other than discharge rate such as changes in discharge patterns, the emergence of burst discharges, or oscillatory activity. Interestingly, STN lesions abolish the increase in GAD mRNA in GPe in par-
exposed to 80 mM K

GABA levels by sample number in the two monkeys. Per our microdialysis protocol, sample 6 was before (black columns) and after (gray columns) treatment with MPTP.

Figure 3. GABA concentrations in the STN as measured with microdialysis in monkeys H and I, before (black columns) and after (gray columns) treatment with MPTP. A and B show the average GABA levels by sample number in the two monkeys. Per our microdialysis protocol, sample 6 was exposed to 80 mw K⁺, resulting in a significant increase in the GABA level. C depicts the average absolute baseline GABA levels in the two monkeys. *p < 0.05 versus the baseline level in the normal state.

GABA activity changes and the development of parkinsonian motor signs

The current rate-based model of the pathophysiology of parkinsonism states that a reduction in GPe discharge leads to the development of the parkinsonian motor abnormalities via disinhibition of STN. This prediction is not supported by the results of our experiments. In contrast to the obvious MPTP-induced parkinsonism, large GPe lesions in normal animals, which, at least in two animals, involved substantial portions of the ventral pallidal motor territory, did not induce discernible motor signs of parkinsonism. These findings are similar to previous studies of smaller GPe lesions (Mink and Thach, 1991), which induced only a slight flexion bias and bradykinesia. As expected, the lesions in our monkeys induced changes in discharge rates in STN and GPi, which were comparable with those seen in the parkinsonian animals. However, GPe lesioned and MPTP-lesioned animals differed with regard to the induction of burst discharges and oscillatory discharge in the basal ganglia. Both parameters were greatly altered in the MPTP-treated animals but were not or only slightly affected in the GPe-lesioned monkeys.

Together, these studies suggest that a reduction in GPe activity and the resulting increases in STN and GPi activity may be necessary but are not sufficient to induce parkinsonism. The findings also suggest that changes in neuronal discharge patterns in the basal ganglia output nuclei may be as relevant as changes in discharge rate in the development of parkinsonian motor signs (Filion, 1979; Bergman et al., 1994; Hassani et al., 1996; Wichmann et al., 1999; Ni et al., 2000; Raz et al., 2000; Bevan et al., 2002; Wichmann and DeLong, 2003a). Even very early studies of the changes in neuronal discharge in the basal ganglia have remarked on the prominence of burst patterns and oscillatory discharge abnormalities in the basal ganglia of parkinsonian monkeys (Filion, 1979; Miller and DeLong, 1987; Filion and Tremblay, 1991; Bergman et al., 1994). Our results are in general agreement with results obtained in rodents in which pallidal lesions and dopamine depletion were also shown to differ (Hassani et al., 1996), although these recordings were performed in anesthetized animals, and behavioral comparisons between the experimental groups were not provided.

It is important to realize that the findings in the GPe-lesioned animals do not rule out the possibility that altered activity in GPe is involved in the development of parkinsonian motor abnormalities. Despite the fact that GPe ablation does not appear to result in the pattern abnormalities or motor abnormalities typical for parkinsonism, less drastic alterations of GPe activity may be related to the induction of burst discharges and oscillatory discharge in the basal ganglia. Both parameters were greatly altered in the MPTP-treated animals but were not or only slightly affected in the GPe-lesioned monkeys.

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<table>
<thead>
<tr>
<th>Monkey</th>
<th>Nucleus</th>
<th>Condition</th>
<th>Average discharge rate (spikes/sec)</th>
<th>Proportion of spikes in bursts (%)</th>
<th>Spectral content (%) 3–8 Hz</th>
<th>Proportion (%) of cells with oscillatory autocorrelogram 3–8 Hz</th>
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<td>H</td>
<td>GPe</td>
<td>Control</td>
<td>65.3 ± 4.5</td>
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<td>35.2 ± 2.3a</td>
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<td>15.72 ± 0.87a</td>
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<td>GPi</td>
<td>Control</td>
<td>65.4 ± 4.3</td>
<td>0.57 ± 0.06</td>
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<td>80.4 ± 4.2a</td>
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<td>17.20 ± 1.46a</td>
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<td>I</td>
<td>GPe</td>
<td>Control</td>
<td>64.8 ± 3.5</td>
<td>0.34 ± 0.06</td>
<td>8.14 ± 0.37</td>
<td>0.05 versa.</td>
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<td>69.5 ± 3.5</td>
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*p < 0.05 against individual control.
synchronized rhythmic bursts in both structures in dopamine-free coculture experiments. The occurrence and maintenance of such bursts may depend on, at least in part, the propensity of STN neurons to respond with rebound bursts to incoming synchronized volleyes of activity along the GPe-STN pathway (Bevan et al., 2002). In favor of a role of GPe in the generation of oscillatory activity in STN and GPi is the fact that the proportion of cells engaged in oscillatory activity in GPe-lesioned animals appeared to be lower than in unlesioned animals (Fig. 2).

Alternatively, oscillatory and burst activity in STN and GPi may emerge independent of GPe input. For instance, it has been speculated that pattern abnormalities in STN may result from loss of dopamine in the STN (Meibach and Katzman, 1979; Campbell et al., 1985; Besson et al., 1988; Johnson et al., 1994; Kreiss et al., 1997; Francois et al., 2000; Smith and Kieval, 2000; Zhu et al., 2002) or from changes in the synchronization between STN neurons and cortex (Magill et al., 2000a,b, 2001; Wichmann et al., 2001b). Other explanations would include altered interactions between STN and brainstem areas such as the pedunculopontine nucleus (Orieux et al., 2000) or between STN and the intralaminar nuclei of the thalamus (Bacci et al., 2002).
Conclusions
Our experiments provide evidence that the neuronal activity in GPe is reduced in the parkinsonian state but do not support the concept that this reduction per se results in parkinsonism. The experiments emphasize the importance of changes in discharge patterns in the basal ganglia output nuclei for the development of parkinsonian motor signs. Although the importance of such changes is obvious, it remains to be seen whether network interactions between STN and GPe, cortex, brainstem or thalamus, or changes in the local milieu within STN are most relevant for the development of pattern changes. Perhaps even more importantly, it remains to be determined which pattern abnormalities are most deleterious for motor performance in parkinsonism.

References


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